

# Upper Arm Anthropometry Is Not a Valid Predictor of Regional Body Composition in Preterm Infants

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## Key Words

Body composition • Magnetic resonance imaging • Preterm infant • Upper arm anthropometry

## Abstract

**Background:** Upper arm anthropometry has been used in the nutritional assessment of small infants, but it has not yet been validated as a predictor of regional body composition in this population. **Objective:** Validation of measured and derived upper arm anthropometry as a predictor of arm fat and fat-free compartments in preterm infants. **Methods:** Upper arm anthropometry, including the upper arm cross-sectional areas, was compared individually or in combination with other anthropometric measurements, with the cross-sectional arm areas measured by magnetic resonance imaging, in a cohort of consecutive preterm appropriate-for-gestational-age neonates, just before discharge. **Results:** Thirty infants born with (mean  $\pm$  SD) a gestational age of  $30.7 \pm 1.9$  weeks and birth weight of  $1,380 \pm 325$  g, were assessed at  $35.4 \pm 1.1$  weeks of corrected gestational age, weighing  $1,785 \pm 93$  g. None of the anthropometric measurements are reliable predictors ( $r^2 < 0.56$ ) of the measurements obtained by magnetic resonance imaging, individually or in combination with other anthropometric measurements. **Conclusion:** Both measured anthropometry and derived upper arm anthropometry are inaccurate predictors of regional body composition in preterm appropriate-for-gestational-age infants.

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## Introduction

Mid-arm circumference (MAC) and skinfold thickness of the arm have been used as simple, noninvasive and inexpensive tools for the assessment of the nutritional status of neonates [1]. Upper arm cross-sectional areas, derived from MAC and triceps skinfold thickness (TS), has been preferred for the assessment of body composition and nutritional status in small infants [2–5], assuming that it would represent a better indicator of the relative contribution of fat and muscle to the total arm area than TS and MAC by themselves [6, 7]. Theoretical assumptions for the calculation of arm muscle and fat areas are based on the fact that the mid-arm is cylindrical, the subcutaneous fat is a concentric ring evenly distributed around a circular core of muscle, the fat thickness is half the TS, and the muscle includes the humeral diameter [6]. Using the equations proposed by Rolland-Cachera et al. [8], it is also assumed that the arm muscle area constitutes a circle surrounded by a rim (fat), the unrolled rim is a rectangle, and the outer circumference of the rim is equal to the inner circumference. Reference values for upper arm cross-sectional areas have been proposed for full-term and preterm neonates [9–11].

Using dual X-ray absorptiometry as a reference method, upper arm anthropometry has been recently validated as proxy for regional and total body composition in neonates [12, 13]. In this age group, the upper arm an-

thropometry shows a good predictive value for total body composition [13], and the TS may give a rough estimate of total body fat [12].

If these were to be consistent, the regional anthropometric measurements capable of predicting total body composition should also be reliable predictors of regional body composition. Upper arm cross-sectional areas measured by anthropometry showed to be good predictors of arm fat and fat-free compartments in adults [14] and children aged 9–15 years [8], using computerized tomography scan or magnetic resonance imaging (MRI) as gold standard methods. In these studies, constants were found to improve the accuracy of anthropometry as predictors of body composition. Validations using gold standard imaging methods of the arm have not yet been performed in neonates. Meanwhile, in several studies the upper arm cross-sectional areas have been used to assess the nutritional status and body composition of small infants [2–5]. However, it is not clear whether the above-mentioned geometrical assumptions used in the calculation of the arm compartments [6, 8] are reliable in this age group. Due to the convenience of bedside measurement of upper arm anthropometry in nutritional assessment, there is a need to validate this method in neonates [15].

The MRI method has been validated as a direct, reliable and noninvasive method for measuring the adipose tissue, capable of quantifying even discrete adipose tissue depots [16–19]. Recently, MRI has been used in the assessment of body composition and nutritional status in small infants [20–22].

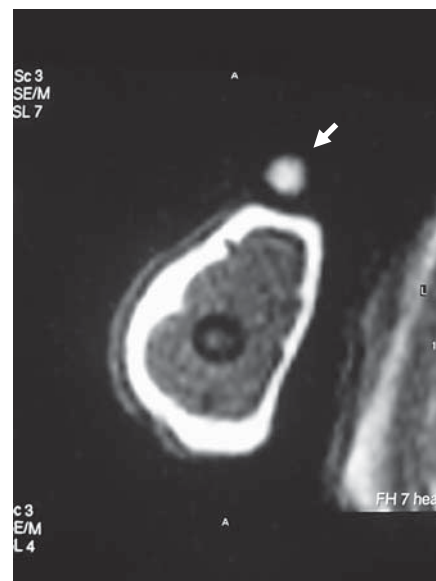
This study aims to validate measured (MAC and TS) and derived upper arm (cross-sectional arm areas) anthropometry as a predictor of regional body fat and fat-free mass in preterm infants, using MRI as the gold standard for measuring the arm compartments.

## Methods

Consecutive newborn infants with gestational age  $\leq 33$  weeks, appropriate for gestational age, were evaluated just before discharge, after obtaining parental consent, and in accordance with the Hospital Ethics Committee. Infants with major congenital malformations, or requiring diuretics or supplemental oxygen at the time of evaluation were excluded.

### Anthropometry

All the anthropometric measurements were performed by the same experienced observer (L.P.d.S.). Body weight (W), length (L), and head circumference (HC) were measured as recommended [23], and MAC and TS were measured as previously described [24]. Indices based on W and L were calculated: the weight-to-



**Fig. 1.** MR image: the hyper-signal of lecithin beacons the most appropriate transaxial image.

length index ( $W/L$ ) =  $W$  (in g)/ $L$  (in cm), the adiposity index ( $AI$ ) =  $W$  (in kg)/ $L^2$  (in m), and the ponderal index ( $PI$ ) =  $100 \times W$  (in g)/ $L^3$  (in cm). The cross-sectional arm areas, such as total arm area (AA), arm muscle area (AMA), and arm fat area (AFA), were calculated using two alternative methods:

Method of Jelliffe and Jelliffe [6]:  $AA = MAC^2/4\pi$ ;  $AMA^J = (MAC - \pi TS)/4\pi$ ;  $AFA^J = AA - AMA^J$ .

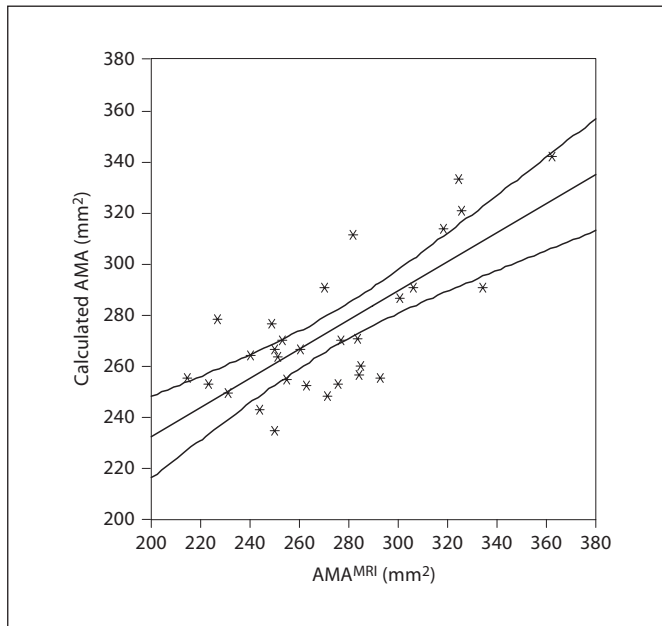
Method of Rolland-Cachera et al. [8]:  $AFA^R = MAC \times TS/2$ ;  $AMA^R = AA - AFA^R$ .

### MRI and Image Analysis

Magnetic resonance images ( $AA^{MRI}$ ,  $AFA^{MRI}$  and  $AMA^{MRI}$ ) were acquired on a 1.5-tesla Philips Gyroscan ACS-NT, Power-Track 1000 scanner. The MRI sequence used in this study was adapted from previously developed scanning protocols involving small infants [20, 21]: a rapid  $T_1$ -weighted spin-echo image sequence, with repetition time = 284 ms, echo time = 18 ms, field of view = 150 mm, number of signal averages = 2, and a  $256 \times 256$  matrix after phase conjugate symmetry. Nine 3-mm-thickness transverse images with 0.3-mm interslice distance were taken during a total acquisition time of 1.58 min.

The infants were positioned supine into a knee coil, and no sedation was used. Considering the  $T_1$  hyper-signal of lecithin [25, 26], a small capsule of lecithin was fixed on the anterior part of the arm, at the previously marked level where the anthropometric measurements were performed. Care was taken to reduce the deformation of soft tissues. The hyper-signal of lecithin permitted to choose the more appropriate transaxial image at the level of the anthropometric measurements (fig. 1).

The same observer (F.A.) defined the criteria for discriminating the subcutaneous adipose tissue (white appearance) from muscle (gray appearance), and manually traced the circumferences: arm ( $MAC^{MRI}$ ), muscle ( $MAMC^{MRI}$ ) and bone. The areas



**Fig. 2.** Validation of the alternative equation for estimation of AMA:  $AMA^{MRI} = (W \times 0.17) + (MAC \times 5.2) - (L \times 6) - 150$ ; W expressed in g, L in cm, and MAC in mm. Coefficient of determination  $r^2 = 0.562$ . Regression line with 95% confidence interval.

included in the circumferences were automatically calculated by pixel counting by the incorporated software.

In pilot measurements, the precision of bone area discrimination was not satisfactory, with an average coefficient of variation (CV) 5.9% (range 1.2–13.8%). Therefore, bone was included in  $AMA^{MRI}$ . The average CV of  $AMA^{MRI}$  was 1.7% (range 0.3–4.7%). The  $AFA^{MRI}$  was obtained by subtracting the  $AA^{MRI}$  from  $AMA^{MRI}$ . The MAC was, therefore, the only direct anthropometric measurement possible to validate by MRI. In the first 20 cases, manual tracing was carried out in triplicate to determine the intraobserver CV. In the last 10 cases, three consecutive acquisitions of images were done to determine the intraobserver CV for the acquisition of images by the same operator.

#### Statistical Analysis

Normal distribution of every variable was verified graphically and through the Kolmogorov-Smirnov test. MRI measurements were correlated individually with each upper arm anthropometry measurement (MAC, TS, and upper arm cross-sectional areas) using the Pearson bivariate correlation test. The predictive value of MRI measurements by each anthropometric measurement and their derived indices was assessed by the coefficients of determination, and was considered as unfit if below 0.5, mediocre if between 0.5 and 0.75, acceptable if between 0.76 and 0.85, good if between 0.86 and 0.95, and excellent if above 0.95.

Multiple regression analysis (enter method) was used to find associations of anthropometric measurements (MAC, TS, AMA, AFA, W, L, W/L, AI, PI and HC) that could be transformed into alternative equations to be correlated with MRI measurements.

**Table 1.** Distribution of the anthropometric measurements

Measurement	Median	Maximum	Minimum	Mean $\pm$ SD
W, g	1,757.5	1,990.0	1,640.0	1,784.7 $\pm$ 93.3
L, cm	43.0	45.2	41.0	43.1 $\pm$ 1.1
HC, cm	30.7	33.0	28.0	30.6 $\pm$ 1.1
MAC, mm	72.2	79.7	67.7	72.9 $\pm$ 0.7
TS, mm	2.5	3.1	2.0	2.5 $\pm$ 0.1
W/L	41.0	46.1	39.1	41.4 $\pm$ 1.9
AI	9.5	10.9	8.8	9.6 $\pm$ 0.5
PI	2.2	2.6	1.9	2.2 $\pm$ 0.2
AA, mm <sup>2</sup>	414.5	505.1	364.4	423.7 $\pm$ 8.0
AMA <sup>I</sup> , mm <sup>2</sup>	328.6	410.9	285.7	337.7 $\pm$ 6.8
AMA <sup>R</sup> , mm <sup>2</sup>	324.0	407.2	280.9	332.8 $\pm$ 6.8
AFA <sup>I</sup> , mm <sup>2</sup>	87.0	111.2	66.2	86.0 $\pm$ 2.6
AFA <sup>R</sup> , mm <sup>2</sup>	91.8	118.2	69.4	90.9 $\pm$ 2.9

**Table 2.** Distribution of the MRI measurements

Measurement	Median	Maximum	Minimum	Mean $\pm$ SD
MAC <sup>MRI</sup> , mm	78.9	92.0	70.2	79.3 $\pm$ 5.1
MAMC <sup>MRI</sup> , mm	60.8	73.1	53.8	61.1 $\pm$ 3.6
AA <sup>MRI</sup> , mm <sup>2</sup>	425.6	540.0	340.5	427.9 $\pm$ 6.5
AMA <sup>MRI</sup> , mm <sup>2</sup>	270.9	362.3	214.7	273.5 $\pm$ 4.7
AFA <sup>MRI</sup> , mm <sup>2</sup>	149.2	227.4	96.4	154.4 $\pm$ 9.06

The validity of the equations was assessed by their coefficient of determination.

Statistical analysis was performed with Microsoft Excel 2000™ and SPSS™ 6.13 (SPSS, Inc., Chicago, Ill., USA) statistical packages.

## Results

Thirty infants born with a (mean  $\pm$  SD) gestational age of  $30.7 \pm 1.9$  weeks and birth weight of  $1,380 \pm 325$  g were evaluated at a corrected gestational age of  $35.4 \pm 1.1$  weeks, when they weighted  $1,786 \pm 93$  g.

The intraobserver CV for the acquisition of images by the operator was 2.0% (0.3–4.8) for  $AMA^{MRI}$ , and 4.4% (0.5–11.4) for  $AFA^{MRI}$ . The intraobserver CV for the manual tracing by the observer was 1.6% (0.6–2.9) for  $AMA^{MRI}$ , and 4.2% (1.6–12.0) for  $AFA^{MRI}$ . No significant differences in MRI measurements between sexes were found.

Body and arm anthropometry (table 1) and measurements obtained by MRI (table 2) are expressed as mean and standard deviation. The MAC proved to be poor predictor ( $r^2 = 0.41$ ), underestimating the  $MAC^{MRI}$  (fig. 2). None of the individual (table 3) or combined (table 4) anthropometric measurements proved to be an acceptable predictor of the MRI measurements:  $r^2 < 0.45$  and  $r^2 < 0.56$ , respectively.

## Discussion

Body weight is commonly used as an isolated measure in the assessment of the nutritional status of small infants in most clinical settings. However, it gives a poor insight into body composition; therefore, weight gain does not necessarily mean growth [27]. The complementary use of some other validated anthropometric measurement predicting regional or total body composition in small infants would represent a useful tool in clinical practice.

Upper arm anthropometry is convenient, particularly in infants that require minimal handling. For this reason, it has been frequently used as a predictor of regional or global body composition in small infants [1–5]. In this population, Schmelzle and Fusch [12] and Koo et al. [13] found a reasonable correlation between the skinfold measurements, including TS, and the total body fat. Others have reported a poor correlation between skinfold measurements and total body fat in newborn infants [22, 27]. This may be due to several factors, including the influence of hydration on the skinfold compressibility [22], and the possibility that skinfold measurements may not reflect nonsubcutaneous fat [21]. In small infants, an unsatisfactory correlation has been described between MAC measurements and body fat and fat-free mass [28], but others have found that MAC measurements may have a reasonable predictive value [13]. Taking into account the contradictory predictive values of TS and MAC found in small infants, validation of upper arm cross-sectional areas assumes special relevance in this population, since their calculation relies on TS and MAC measurements.

Compared with ultrasound measurements, upper arm cross-sectional areas derived by anthropometry may lead to overestimation of muscle and underestimation of fat in full-term neonates [24]. In this population, Kabir and Forsum [29] also found that TS underestimates the adipose tissue in comparison with ultrasound measurements. Like anthropometry, the ultrasonic method has shortcomings when measuring subcutaneous tissue [30] and cross-sectional muscle areas, at least in adults [31].

**Table 3.** Coefficient of determination ( $r^2$ ) of the individual anthropometric measurements

Anthropometry	MRI		
	$AA^{MRI}$ ( $427.9 \pm 6.5$ $mm^2$ )	$AMA^{MRI}$ ( $273.5 \pm 4.7$ $mm^2$ )	$AFA^{MRI}$ ( $154.4 \pm 9.1$ $mm^2$ )
W ( $1,784.6 \pm 93.3$ g)	–	0.372	0.024
L ( $47.1 \pm 1.1$ cm)	–	0.032	0.020
HC ( $30.6 \pm 1.1$ cm)	–	0.012	0.020
MAC ( $72.9 \pm 0.7$ mm)	–	0.449	0.048
TS ( $2.5 \pm 0.1$ mm)	–	–	0.058
W/L ( $41.4 \pm 1.9$ )	–	–	0.010
AI ( $1.0 \pm 0.05$ )	–	–	0.000
PI ( $2.2 \pm 0.2$ )	–	–	0.001
AA ( $423.7 \pm 8.0$ $mm^2$ )	0.422	0.449	0.048
AFA <sup>J</sup> ( $86.0 \pm 2.6$ $mm^2$ )	–	–	0.410
AFA <sup>R</sup> ( $90.9 \pm 2.9$ $mm^2$ )	–	–	0.397
AMA <sup>J</sup> ( $337.7 \pm 6.8$ $mm^2$ )	–	0.073	–
AMA <sup>R</sup> ( $332.8 \pm 6.7$ $mm^2$ )	–	0.073	–

**Table 4.** Coefficient of determination ( $r^2$ ) of the combined anthropometric measurements, including anthropometric indexes

Anthropometry	MRI	
	$AMA^{MRI}$	$AFA^{MRI}$
MAC and L	0.449	–
MAC and W	0.533	0.053
MAC, W and L	0.562	–
MAC, W, L and HC	0.562	–
TS and W	–	0.068
TS, W and HC	–	0.068
AMA <sup>J</sup> and L	0.410	–
AMA <sup>J</sup> and W	0.518	–
AMA <sup>J</sup> , W and L	0.548	–
AMA <sup>J</sup> , W, L and HC	0.548	–
AMA <sup>R</sup> and L	0.410	–
AMA <sup>R</sup> and W	0.504	–
AMA <sup>R</sup> , W and L	0.548	–
AMA <sup>R</sup> , W, L and HC	0.548	–
AFA <sup>J</sup> and W	–	0.078
AFA <sup>J</sup> , W and HC	–	0.116
AFA <sup>R</sup> and W	–	0.078
AFA <sup>R</sup> , W and HC	–	0.116
AFA <sup>J</sup> and W/L	–	0.075
AFA <sup>J</sup> and AI	–	0.078
AFA <sup>J</sup> and PI	–	0.081
AFA <sup>R</sup> and W/L	–	0.074
AFA <sup>R</sup> and AI	–	0.077
AFA <sup>R</sup> and PI	–	0.081



In this study, the MRI was chosen as the gold standard method for measuring the arm compartments. This direct and noninvasive method has been validated in animals [17], human cadavers [18], and human adults [19] for the assessment of adipose tissue. It has also been used in lean tissue assessment in human beings, although with lower precision [32]. The excellent precision in identifying the adipose tissue [16] has been important for distinguishing subcutaneous adipose tissue from muscle, permitting a more reliable manual tracing of the arm circumference and the inner limit of the subcutaneous fat ring separating subcutaneous adipose tissue from muscle (muscle circumference). Considering the poor precision of bone MRI measurements in this study, bone was included in AMA<sup>MRI</sup>, comparing reasonably well to anthropometric AMA, which also includes bone area.

The MRI measurements were correlated with upper arm anthropometric measurements both individually and in combination with other anthropometric measurements. Among the individual measurements, MAC had a surprisingly poor predictive value. In very premature infants, this may be explained by the difficulty in adjusting perfectly the tape around their arms with very soft tissues, leading to a significant compression of the skin. To avoid compression, small spaces may be left between the wrinkled skin. By contrast, the MAC measured by MRI is not affected by compression, and its reliability is mainly dependent on the observer's manual tracing.

The rationale for combining anthropometric measurements was to join on the same equation several anthropometric measurements that might reflect the same body compartments, a methodology used by others [13]. For instance, L was combined with AMA derived by anthropometry when correlated with AMA<sup>MRI</sup>, since L is considered a global indicator of lean body mass [33]; indices based on the W and the L, such as PI [21], AI [34] and W/L [35] may roughly reflect the adiposity. Both ap-

proaches failed to find any acceptable predictor of the upper arm compartments.

The present results are not fully consistent with the results of Schmelzle and Fusch [12] and Koo et al. [13], probably due to some methodological differences. Both studies [12, 13] evaluated the predictive value of regional and whole body anthropometry for total body composition using dual X-ray absorptiometry as the gold standard, while we considered MRI as more adequate for morphological measurements and estimation of regional body composition. The evaluation of upper arm anthropometry as predictor of whole-body composition was beyond the scope of our study. Both studies [12, 13] analyzed mixed samples of full-term and preterm neonates, including appropriate for gestational age, large and small for gestational age neonates, assessed during the first days of life; the patients in the study of Schmelzle and Fusch [12] were also assessed at 2 and 4 months of age. Our study focused only on appropriate-for-gestational-age preterm babies, assessed at discharge.

Our results confirm that both measured anthropometry and derived upper arm anthropometry, either isolated or in combination with the other anthropometric measurements, are inaccurate predictors of regional body composition in appropriate-for-gestational-age preterm infants. This raises concerns about the validity of previous studies in this population assessing the body composition and nutritional status based on upper arm anthropometry; therefore, the results of these studies should be interpreted with caution.

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